(19)日本国特許庁 (JP)

(12) 公開特許公報(A)

(11)特許出願公開番号

特開平5-76762

(43)公開日 平成5年(1993)3月30日

(51)Int.Cl. ⁵ B 0 1 J 23/84 B 0 1 D 53/36 B 0 1 J 23/34 23/76 23/78	102 B A A	庁内整理番号 8017-4G 9042-4D 8017-4G 8017-4G 8017-4G	FΙ	技術表示箇所
			審査請求 未請求	: 請求項の数 1(全 12 頁) 最終頁に続く
(21)出願番号	特顏平3-315589 平成3年(1991)9	月21日	(71)出願人 (72)発明者 (72)発明者 (72)発明者	界化学工業株式会社 大阪府堺市戎之町西1丁1番23号 仲辻 忠夫 大阪府堺市戎島町5丁1番地 堺化学工業 株式会社中央研究所内 奥野 雅雄 大阪府堺市戎島町5丁1番地 堺化学工業 株式会社中央研究所内

(54) 【発明の名称 】 窒素酸化物接触還元用触媒

(57)【要約】

【構成】下記(1)式で表されるペロブスカイト型複合酸化物が固体酸担体に担持されてなる。

Ax B1 - x Cy C1 - y O3

(1)

(式中、AはLaまたはCe、BはBa、Sr、Ca、Mg、Pb、ZnまたはAg、CはMn又はCo、CはFe、Ni、Cr、Cu、V、Mo、W、Ti、Zr、Nb、Pd、Rh、RuまたはPtであり、また $0 \le X \le 1$ 、 $0 \le Y \le 1$ である。)

【効果】工場、自動車などから排出される排ガスの中に 含まれる有害な窒素酸化物、COおよび炭化水素を広い A/Fの範囲において効率良く浄化することができる。

【特許請求の範囲】

【請求項1】下記(1)式で表されるペロブスカイト型 複合酸化物が固体酸担体に担持されてなることを特徴と する排ガス中の窒素酸化物、炭化水素化合物及び一酸化 炭素を同時に除去するための触媒。

Ax B_{1 - x} C_y C_{1 - y} O₃ (1) (式中、AはLaまたはCe、BはBa、Sr、Ca、 Mg、Pb、Zn又はAg、CはMn又はCo、CはF e, Ni, Cr, Cu, V, Mo, W, Ti, Zr, N b、Pd、Rh、Ru又はPtであり、また0≦X≦ $1, 0 \le Y \le 1$ \forall $0 \le Y \le 1$

【発明の詳細な説明】

【産業上の利用分野】本発明は窒素酸化物接触還元用触 媒に係わり、詳しくは工場、自動車などから排出される 排ガスの中に含まれる有害な窒素酸化物、炭化水素及び 一酸化炭素を同時に除去する際に使用する三元触媒に関 する。

【従来の技術及び発明が解決しようとする課題】排ガス 中のNOx、HC及びCOなどの有害成分を同時に除去 する反応はNOxをNaに還元する反応とHC及びCO 20 を二酸化炭素と水に酸化する反応が同時に進行する反応 であるため、排ガス中に含まれる酸素、NOx などの酸 化性成分とHC、COなどの還元性成分の量比が極めて 重要となるので、理論空燃比に近い条件で排ガス処理が 行われている。しかし、理論空燃比での運転は燃費の悪 化の一因となり、ウインドーの広い触媒の開発が望まれ てきた。従来三元触媒成分として、白金ーロジウム、バ ラジウム-ロジウム、白金-パラジウム-ロジウム触媒 が高い活性を示し、アルミナに白金-ロジウムを担持し た触媒が実用化されている。また、これらの貴金属触媒 30 は高価であるためこれに代わる触媒としてベロブスカイ ト型複合酸化物あるいはこれらにパラジウムを含有させ る触媒などが提案されてきた。しかし、これらの触媒の いづれもが理論空燃比あるいはその近辺比でのみ高い活 性を示した。(ウィンドウの幅が狭い)そのため燃費を 犠牲にすることなく幅の広いウィンドウを有する触媒の 開発が望まれてきた。本発明は、以上の事情に鑑みなさ れたものであって、その目的とするところは、幅の広い ウィンドウを有する三元触媒、言い換えれば理論空燃比*

*からのずれが大きい排ガス条件下においても有効に機能 する三元触媒を提供することにある。

【課題を解決するための手段】本発明に係る炭化水素及 び/又は含酸素化合物を還元剤として使用する窒素酸化 物接触還元用触媒は、下記(2)式で表されるペロブス カイト型複合酸化物が、ゼオライト、結晶性珪酸燐酸ア ルミニウム (SAPO)、結晶性燐酸アルミニウム (A LPO)、結晶性燐酸金属アルミニウム(MAPO)、 アルミナ、チタニア、ジルコニア、シリカーアルミナ等 10 の固体酸担体に担持されてなる。

Ax Bı - x Cy Cı - y Os (式中、AはLa又はCe、BはBa、Sr、Ca、M g、Pb、Zn又はAg、CはMn又はCo、CはF e, Ni, Cr, Cu, V, Mo, W, Ti, Zr, N b、Pd、Rh、RuまたはPtであり、また0≦X≦ 1、0≦Y≦1である。) 本発明における固体酸担体と は、使用される温度領域において固体酸性を示す担体を いう。固体酸性の確認は、アンモニアを用いた昇温脱離 法、あるいはアンモニア又はピリジンを用いたin s itu FTIR法によりなされる。固体酸担体として は、次に示すゼオライト系固体酸担体や酸化物系固体酸 担体などがある。

(i)ゼオライト系固体酸担体は、Na-モルデナイ ト、Na-ZSM-5、Na-USY (USY:ウルト ラステイブルY型ゼオライト)等の耐熱性に優れるゼオ ライトを硫酸アンモニウム等のアンモニウム塩の水溶液 又は硫酸等の酸で処理して、ゼオライト中のアルカリ金 属の一部又は全部をアンモニウムイオン(NH⁺ +)又 は水素イオン(H⁺)にイオン交換することにより得ら れる。NH⁴ [†] でイオン交換する方法による場合は、最 後に焼成処理を必要とする。ゼオライト系固体酸担体と しては、例えば下記(3)式で表されるモルデナイト型 ゼオライトを酸処理して得られる酸型モルデナイトであ って、SiO2/Al2O3のモル比が13~20であ り、且つ、SiO₂ /Al₂O₃ のモル比が25~20 0である酸型モルデナイトや、下記(4)式で表される ゼオライト中のイオンMの一部又は全部をTi^^、 Z r 4 + 又はSn 4 + でイオン交換して得られるゼオライ トが挙げられる。

 M_2 [(AlO₂)₂, (SiO₂)₁₀] · ZH₂O

(3)

(ただし、式中、Mはアルカリ金属イオン、rはゼオラ※ ※イトの合成条件により変動する値である。) M'_{A} [(AlO₂), (SiO₂),] · Z' H₂ O

4)

(ただし、式中、イオンM'はアルカリ金属イオン、ア ルカリ土類金属イオン又は水素イオン、nA=p(n は、イオンMの価数である)、q/p≥5である。) (ii)酸化物系固体酸担体としては、Al2〇。、T $i O_2$ 、 $T i O_2$ / $S O_4$ - - 、 $Z r O_2$ 、 $Z r O_2$ / S〇4 - 等の単一金属酸化物や、SiO2 / Al2 〇 50 似の多孔構造又は層状構造を有する一種の結晶性燐酸ア

s、TiO2/Al2Os、TiO2/ZrO2等の複 合酸化物等が挙げられる。これらの中では、耐熱性の点 で、Al2Os、ZrO2、SiO2/Al2Osが好 ましい。

(i i i) 固体酸担体の他の例としては、ゼオライト類

3

ルミニウム (ALPO) や、その近縁物質である結晶性 珪酸燐酸アルミニウム(SAPO)、ALPOの燐又は 燐-アルミニウムの一部をチタン、鉄、マグネシウム、 亜鉛、マンガン、コバルト等の金属で置換した結晶性燐 酸金属アルミニウム(MAPO)などが挙げられる。A LPOタイプの燐酸塩は、上記の燐酸源及び金属源と、 シリカ、シリカゾル、珪酸ソーダなどの中から選ばれた 所望の組合せに、アミン、第四級アンモニウム等のいわ ゆるテンプレートを混合した原料から、ゼオライトを合 成する場合と類似した条件下で水熱合成法により調整す ることが出来る。ゼオライトを合成する場合との主な相 違点は、一般により高温(概ね150℃以上)でpH酸 性領域で合成されることである。ALPOタイプの燐酸 塩の組成は、一般にA1203・(0.8~1.2)・ P₂ O₅ · n H₂ Oで表される。また、SAPO又はM APOの場合においては、置換するシリカ及び金属の最 大量は、アルミニウム及び燐の総量の約1/10程度で あるが、本発明においては、必ずしもこの組成範囲に入 っていないもの、すなわち非晶質を含んでいるものを使 用してもよい。水熱合成法により得られるALPOタイ 20 プの燐酸塩を担体として使用する場合は、一般に、水 洗、乾燥した後、空気中で焼成して残存しているテンプ レートを焼却除去したものが使用される。本発明に係る 触媒は、例えば次に示す(1)、(2)又は(3)の方 法により調整することができる。

(1)固体酸担体を分散させたスラリー中に、La、Ce、Ba、Sr、Ca、Mg、Pb、Zn、Ag、Mn、Co、Fe、Ni、Cr、Cu、V、Mo、W、Ti、Zr、Nb、Pd、Rh、Ru又はPtの硝酸塩等の水溶性塩や、これらのアルコキシドにアルコール溶液 30を投入し、これらを中和あるいは加水分解させる方法などによって、固体酸担体にこれらの複合金属種の水酸化物等のペロブスカイト化合物前駆体を担持させる。次いで、濾過、水洗、リバルブを繰り返し行った後、乾燥し、焼成する。

(2)固体酸担体と、別途調整したペロブスカイト化合物とを、遊星ミルなどによって充分に湿式粉砕混合する。

(3)固体酸担体の水溶性塩又は水酸化物等の前駆体と、La、Ce、Ba、Sr、Ca、Mg、Pb、Z 40 n、Ag、Mn、Co、Fe、Ni、Cr、Cu、V、Mo、W、Ti、Zr、Nb、Pd、Rh、Ru又はPtの硝酸塩等の水溶性塩や、これらのアルコキシドのアルコール溶液とを均質に混合した溶液を、中和又は加水分解させる方法などによって沈澱物を生成させる。次いで、この沈澱物を濾過、水洗、リバルブを繰り返し行った後、乾燥し、焼成する。以上の方法において、ペロブスカイト化合物の生成温度は低い方が好ましい。その理由は、その生成温度が低いほど、大きな比表面積を有するペロブスカイト化合物が得られ、固体酸担体と、ペロ 50

ブスカイト化合物を構成する元素との反応により固体酸 担体の固体酸性が変質したり、ペロブスカイト化合物の 生成量の低下により触媒の活性が低下したりすることを 回避することができるからである。したがって、Al2 O_s、TiO₂などのようにペロブスカイト化合物を構 成する元素との反応性が高い固体酸担体を用いる場合に あっては、固体酸担体を構成する元素とペロブスカイト 化合物を構成する元素との均質性を高める(3)の方法 は好ましくない。一般には、(1)の方法が好ましい が、(2)の方法によってもかなり高い活性を示す触媒 を得ることができる。ペロブスカイト化合物の好適な担 持量は、このペロブスカイト化合物と固体酸担体との総 重量に対して、1.0~50重量%である。50重量% を越えると、増量に応じた添加効果が得られないばかり でなく酸素が共存する反応系においては酸素による炭化 水素や一酸化炭素の浄化率が高くなるもののNOxの浄 化率が大きく低下する。一方、1.0重量%未満ではN O、炭化水素、COのいづれの浄化率をも充分向上させ ることができない。本発明に係る触媒は、従来公知の成 形方法によりハニカム状、球状等の種々の形状に成形す ることができる。この成形の際に、成形助剤、成形体補 強体、無機繊維、有機パインダーなどを適宜配合しても よい。また、予め成形された基材上にウォッシュコート 法などにより、被覆担持させることもできる。その他、 従来公知の触媒調整法によることもできる。本発明に係 る触媒が窒素酸化物、炭化水素及び一酸化炭素に対して 浄化活性を示す最適な温度は、触媒種により異なるが、 通常100~800℃であり、この温度領域において は、空間速度(SV)500~100,000程度で排 ガスを通流させることが好ましい。なお、より好適な使 用温度領域は200~800℃である。また本発明に係 る触媒が有効に機能する空燃比(A/F)は、14.0 ~60であり、好ましくは14.0~40、より好まし くは14.0~30である。

【実施例】以下、本発明を実施例に基づいて更に詳細に 説明するが、本発明は下記実施例に何ら限定されるもの ではなく、その要旨を変更しない範囲において適宜変更 して実施することが可能なものである。

(1)触媒の調整

40 実施例1

La (NO₃)₂・6H₂O、Mn (Ac)₂・4H₂O (Ac:CH₃COOH、以下も同じ)、Sr (NO₃)₂、Co (NO₃)₂・6H₂Oを、各々101.05g、28.60g、74.08g、135.83g 秤量し、500mlの水に溶かした。この溶液に、充分に攪拌しながら121g/1のNaOH水溶液を加えて、pHを10とした。中和反応終了後、18時間攪拌を続け熟成を行った。その後、濾過、水洗、リバルブを、濾過水の導電率がリバルブ用水とほぼ同じになるまで繰り返した。得られた濾過ケーキを120℃で18時

間乾燥し、次いで700℃で3時間焼成した。得られた 焼成物のXRDを求めた結果、ペロブスカイト結晶相が 生成していることが判った。また、この焼成物のBET 法による比表面積(以下の比表面積も同法による)は、 23. 7m²/gであった(Lao, 4 Sro, 6 Co 。. a Mno. 2 Os)。 とのようにして得たペロプス カイト化合物30gと、硫酸法酸化チタン工程により得 たメタチタン酸 (Ti〇2 · H2 O) を600℃で3時 間焼成して得た活性酸化チタン(比表面積:104.2 m² /g) 100gとの混合物に、水100gを加え、 遊星ミルにて30分間粉砕混合し、更に水で粘度調整し てウォッシュコート用スラリーを得た。このスラリーを 1. 25 mmピッチのコージュライト社製のハニカムに 塗布して触媒を担持させ、試作サンプル(A-1)を得 た。このときのスラリーの塗布量は、ハニカム1cc当 たり0.116gであった。

実施例2

La $(NO_3)_2 \cdot 6H_2 O \geq Mn (Ac)_2 \cdot 4H_2$ Oとを、各々89.5g、50.66g秤量し、次い で、実施例 I と同様の方法にて、ペロブスカイト化合物 20 (LaMnO。)を得た。このペロブスカイト化合物の 比表面積は、29.1m²/gであった。このようにし て得たペロブスカイト化合物30gと、住友化学社製の γ-アルミナ100gとの混合物に、水100g加えて スラリーを得、次いで実施例1と同様のハニカムに塗布 して触媒を担持させ、1.25mmピッチのハニカム形 状の試作サンブル(A-2)を得た。このときのスラリ 一の塗布量は、ハニカム1cc当たり0.094gであ った。

実施例3

La $(NO_3)_2 \cdot 6H_2 O$, Pb $(NO_3)_2$, Mn (Ac) 2 · 4 H 2 Oを、各々7 1. 60g、13. 6 9g、50.66g秤量して混合し、次いで、800℃ で3時間焼成したこと以外は実施例 I と同様の方法に て、ペロブスカイト化合物(Lao. a Pbo. 2 Mn 〇。)を得た。このペロブスカイト化合物の比表面積 は、23.7m²/gであった。このようにして得たべ ロブスカイト化合物30gと、住友化学社製のャーアル ミナ100gとの混合物に、水100g加えてスラリー を得、次いで実施例1と同様のハニカムに塗布して触媒 40 を担持させ、1.25mmピッチのハニカム形状の試作 サンブル (A-3) を得た。このときのスラリーの塗布 量は、ハニカム1cc当たり0.117gであった。 実施例4

La $(NO_3)_2 \cdot 6H_2 O$, Co $(NO_3)_2 \cdot 6H$ 2 〇を、各々88.07g、59.19g秤量して混合 し、次いで、800℃で3時焼成したこと以外は実施例 Iと同様の方法にて、ペロブスカイト化合物 (LaCo 〇。)を得た。このペロブスカイト化合物の比表面積

ペロブスカイト化合物30gと、日本化学社製のH形モ ルデナイト(HM-23)100gとの混合物に、水1 00g加えてスラリーを得、次いで実施例1と同様のハ ニカムに塗布して触媒を担持させ、1.25mmピッチ のハニカム形状の試作サンプル(A-4)を得た。この

ときのスラリーの塗布量は、ハニカム1 c c 当たり0.

113gであった。

実施例5

Ce $(NO_3)_2 \cdot 6H_2 O$, Ba $(NO_3)_2$, Co (NO₃)₂・6H₂ Oを、各々70、65g、10. 63g、59.19g秤量し、次いで実施例1と同様の 方法にて、ペロブスカイト化合物(Coo. a Ba a. 2 CoOs)を得た。このペロブスカイト化合物の 比表面積は、23.0 m² /gであった。とのようにし て得たペロブスカイト化合物30gと、日本化学社製の H形モルデナイト(商品名「HM-23」)100gと の混合物に、水100g加えてスラリーを得、次いで実 施例1と同様のハニカムに塗布して触媒を担持させ、 1. 25 mmピッチハニカム形状の試作サンプル (A-5)を得た。このときのスラリーの塗布量はハニカム1 cc当たり0.130gであった。

実施例6

La $(NO_3)_2 \cdot 6H_2 O$, Mn $(Ac)_2 \cdot 4H_2$ O、TiCl。水溶液 (Tiとして14.82g/10 0m1濃度の水溶液)を、各々90.84g、25.7 1g、339.0m1秤量し、次いで、実施例1と同様 の方法にて、ペロブスカイト化合物(LaMno, sT io. oOs)を得た。このペロブスカイト化合物の比 表面積は、25.3m²/gであった。このようにして 得たペロブスカイト化合物10gと、日本アロジル社製 のシリカーアルミナ(商品名「COK-84」)100 gとの混合物に、水100g加えてスラリーを得、次い で実施例1と同様のハニカムに塗布して触媒を担持さ せ、1.25mmピッチのハニカム形状の試作サンプル (A-6)を得た。このときのスラリーの塗布量は、ハ ニカム1 c c当たり0.098gであった。

実施例7

<ペロブスカイト化合物の調製>白水化学工業社製La エトキシドのエタノール溶液(LagOgとして73g /1 濃度の溶液)、同社製のBaエトキシドのエタノー ル溶液(BaOとして79g/1濃度の溶液)、同社製 Niエトキシドのエタノール溶液(NiOとして67g /1濃度の溶液)、Coエトキシドのエタノール溶液 (CoOとして91g/1濃度の溶液)を、各々10 0. 0m1, 21. 74m1, 49. 95m1, 9. 2 2m1秤量して混合し、充分攪拌しながら1重量%のア ンモニア水溶液を徐々に滴下し加水分解させた。次い で、これを均一混合しながら蒸発乾固させ、800℃で 3時間焼成してペロブスカイト化合物 (Lao. Ba は、17.47m²/gであった。このようにして得た 50 o.2 Coo.a Nio.2 Os)を得た。このペロブ

スカイト化合物の比表面積は36.9 m²/gであっ た。

<SAPO-34の調製>水129.6gに、攪拌しな がら細かく砕いたアルミニウムイソプロポキシド90. 7gを少量づつ加え、均一になるまで攪拌混合した。と の混合液に、85%燐酸水溶液51.3gを滴下し、均 一になるまで攪拌混合した後、さらに50%シリカゾル 16.0g加え、充分に攪拌混合した。次いで、水酸化 テトラエチルアンモニウム81.6gを加え、充分に攪 拌混合した。この混合物を、オートクレーブに仕込み、 200℃で24時間攪拌反応させた後、生成物を濾過分 離し、さらに水洗、乾燥した後、500℃で3時間空気 中で焼成してSAPO-34を得た。このSAPO-3 4は、Si、Al、Pをそれぞれ9.5、18.0、1 9. 0重量%含有する組成のものであった。このように して得たペロブスカイト化合物25gと、100gのS APO-34との混合物に、水100gを加えてスラリ ーを得、次いで実施例1と同様のハニカムに塗布して触 媒を担持させ、1.25mmピッチのハニカム形状の試 作サンプル (A-7) を得た。このときのスラリーの塗 20 布量は、ハニカム1cc当たり0.138gであった。 実施例8

La $(NO_3)_3 \cdot 6H_2 O$, Sr $(NO_3)_2$, Co $(NO_3)_2 \cdot 6H_2 O$, Fe $(NO_3)_2 \cdot 6H_2 O$ を、各々101.05g、74.08g、135.83 g、33.60gを秤量して混合し、実施例1と同様の 方法にて、ペロブスカイト化合物(Lac. Sr o. a Coo. a Feo. 2 Os)を得た。このペロブ スカイト化合物の比表面積は、21.6m²/gであっ た。このようにして得たペロブスカイト化合物30g と、住友化学社製のγ-アルミナ100gとの混合物 に、水100g加えてスラリーを得、次いで実施例1と 同様のハニカムに塗布して触媒を担持させ、1.25m mピッチのハニカム形状の試作サンプル(A-8)を得 た。このときのスラリーの塗布量は、ハニカム1 c c 当 たり0.098gであった。

実施例9

La $(NO_3)_3 \cdot 6H_2 O$, Zn $(NO_3)_2 \cdot 6H$ 2 O, Co (NO₃) 2 · 6 H₂ O, Cu (NO₃) 2 ·3H₂ Oを、各々202、10g、28、41g、1 35.83g、28.19gを秤量して混合し、実施例 1と同様の方法にて、ペロブスカイト化合物(La o. a Zno. g Coo, a Cuo. g Os)を得た。 このペロブスカイト化合物の比表面積は、17.3m² /gであった。このようにして得たペロブスカイト化合 物30gと、住友化学社製のγ-アルミナ100gとの 混合物に、水100g加えてスラリーを得、次いで実施 例1と同様のハニカムに塗布して触媒を担持させ、1. 25mmピッチのハニカム形状の試作サンプル(A-

1 c c 当たり0.083 g であった。

実施例10

La (NO₃)₃ · 6 H₂ O, AgNO₃, Co (NO s) 2 · 6 H 2 O、Zr (NOs) 4 · 5 H 2 Oを、各 7202.10g, 19.82g, 135.83g, 6 9.87gを秤量して混合し、実施例1と同様の方法に て、ペロブスカイト化合物 (Lao.s Ago. 2 Co o. a Zro. 2 Os)を得た。このペロプスカイト化 合物の比表面積は、17.3 m²/g であった。このよ うにして得たペロブスカイト化合物30gと、住友化学 社製のγ-アルミナ100gとの混合物に、水100g 加えてスラリーを得、次いで実施例1と同様のハニカム に塗布して触媒を担持させ、1.25mmピッチのハニ カム形状の試作サンブル(A-10)を得た。このとき のスラリーの塗布量は、ハニカム1cc当たり0.09 7gであった。

実施例11

La $(NO_3)_3 \cdot 6H_2 O$, Sr $(NO_3)_2$, Co (NOs) 2 · 6 H 2 O、Cr (NOs) s を、各々1 01. 05g, 74. 08g, 135. 83g, 27. 79gを秤量して混合し、実施例1と同様の方法にて、 ペロプスカイト化合物(Lao. a Sro. a Co o. a Cro. 2 Os)を得た。このペロブスカイト化 合物の比表面積は、20.3m²/gであった。 <ALPO-5の調製>85%燐酸69.2gと水17 8gとの混合物に、擬ベーマイト粉末(アルミナ67 %、酢酸9.5%を含むもの)45.8gを少量づつ加 え、均一になるまで攪拌混合した。この液に、トリプロ ピルアミン43.8gを加え、均一になるまで攪拌混合 30 した。この混合物をオートクレーブに仕込み、150℃ で70時間攪拌反応させた後、生成物を濾過分離し、水 洗、乾燥した後、500℃で3時間空気中で焼成してA LPO-5を得た。このALPO-5は、A1、Pをそ れぞれ18.0、22.0重量%含有する組成のもので あった。このようにして得たペロプスカイト化合物30 gと、100gのALPO-5との混合物に、水100 g加えてスラリーを得、次いで実施例1と同様のハニカ ムに塗布して触媒を担持させ、1.25mmピッチのハ ニカム形状の試作サンブル(A-11)を得た。このと きのスラリーの塗布量は、ハニカム1cc当たり0.1 04gであった。

実施例12

<ペロブスカイト化合物の調製>La(NOs)2・6 $H_2 O_s Sr (NO_3)_2 Co (NO_3)_2 \cdot 6H_2$ O、NbCl。(塩酸水溶液、Nbとして50g/l濃 度の溶液)を、各々101.05g、74.08g、1 35.83g、216.81m1秤量して混合し、以下 実施例1とにして、ペロブスカイト化合物(Lao. 4 Src. a Coc. a Nbc. 2 Os)を得た。このペ 9)を得た。このときのスラリーの塗布量は、ハニカム 50 ロブスカイト化合物の比表面積は、18.9 m²/gで

あった。

<MAPO-5の調製>酢酸第一マンガン4.9gと酢 酸第二銅4.1gとを水129gに溶解した液に攪拌し ながら細かく砕いたアルミニウムイソプロプキシド5 6.3gを少量ずつ加え、均一になるまで攪拌混合し た。この液に、85%燐酸55.4g、ジエチルエタノ ールアミン56.3g、水55.5gの混合液を攪拌し ながら少量ずつ加え、均一になるまで攪拌混合した。と の液をオートクレーブに仕込み、200℃で25時間反 応させた後、生成物を濾過分離し、水洗、乾燥した後、 500℃で3時間空気中で焼成してMAPO-5を得 た。このMAPO-5は、A1、P、Mn、Cuをそれ ぞれ19.0、19.0、2.8、4.4重量%含有す る組成のものであった。このようにして得たペロブスカ イト化合物25gと、100gのMAPO-5との混合 物に、水100g加えてスラリーを得、次いで実施例1 と同様のハニカムに塗布して触媒を担持させ、1.25 mmピッチのハニカム形状の試作サンプル(A-12) を得た。このときのスラリーの塗布量は、ハニカムlc c当たり0.116gであった。

実施例13

実施例1において、活性酸化チタンに代えて水酸化ジル コニウムを600℃で3時間焼成して得た2 r O2 (比 表面積: 148.3 m²/g)を用いたこと以外は実施 例1と同様にして試作サンブル(A-13)を得た。こ のときのスラリーの塗布量は、ハニカム1 c c 当たり 0. 139gであった。

実施例14

<ペロブスカイト化合物の調製>La(NO₃)₂・6 $H_2 O$, $Sr (NO_3)_2$, $Co (Ac)_2 \cdot 4H_2 O$ を、各々88.07g、10.76g、50.66g秤 量し、500mlの水に溶かした。121g/lのNa OH水溶液を充分に攪拌しながら滴下して、液のpHを 10とした。中和反応終了後、18時間攪拌を続け熱成 を行った。その後、濾過、水洗、リバルブを、濾過水の 導電率がリバルブ用水のそれとほぼ同じになるまで繰り 返した後、濾過ケーキを120℃で18時間乾燥した。 この乾燥物を粉砕し、この粉砕物にシュウ酸バナジル溶 液(Vとして100g/1濃度の水溶液) 25.90m 1を加え充分混練した御蒸発乾固し、120℃で18時 40 間乾燥し、次いで850℃で3時間焼成してペロブスカ イト化合物 (Lac. s Sro. z Coo. s Vo. z 〇。)を得た。このペロブスカイト化合物の比表面積 は、12.8m²/gであった。 (シリカージルコニア の調製>

日産化学社製のシリカゾルO型(SiO2として20重 量%濃度のもの)と塩化ジルコニウム(2 r C 1 ₄)を 各々100.0g、97.20g秤量し攪拌しながら充 分混合し、水にて総量を500mlとした。この液に、

した。中和反応終了後、18時間攪拌を続け、その後濾 過、水洗、リパルブを繰り返して濾過ケーキを得、との 瀘過ケーキを120℃で18時間乾燥し、3時間焼成し た。得られた焼成物の比表面積は、297m2/gであ った。このようにして得られたペロブスカイト化合物3 0gと、上記焼成物100gとの混合物に、水100g を加え、遊星ミルにて30分間乾粉砕混合し、さらに水 で粘度調製してウォッシュコート用スラリーを得た。と のスラリーを1.25mmピッチのコージュライト社製 10 のハニカムに塗布して触媒を担持させ、試作サンブル (A-14)を得た。このときのスラリーの塗布量は、 ハニカム1cc当たり0.127gであった。 実施例15

<ペロブスカイト化合物の調製>実施例14において、 シュウ酸パナジル水溶液に代えてモリブデン酸アンモニ ウムのアンモニア性水溶液(MoOg として25g/1 濃度の水溶液) 260.22mlを加えたこと以外は実 施例14と同様にしてペロブスカイト化合物(La o. s Sro. 2 Coo. s Moo. 2 Os)を得た。 20 <Zr-モルデナイトの調製>日本化学社製のNaモル デナイト (NM-100P) 100g を硝酸ジルコニル 水溶液 (ZrO2 として100g/1濃度のもの) に浸 漬し攪伴しながら70℃に1時間保持し、NaをZrと イオン交換させた。濾過、水洗して得たゼオライトケー キを乾燥後650℃で4時間焼成した。このゼオライト (Zr-モルデナイト)のZrの含有量は、3.3重量 %であり、また比表面積は391m²/gであった。と のようにして得たペロブスカイト化合物30gと、Zr -モルデナイト100gとの混合物に、水を100g加 え、遊星ミルにて30分間粉砕混合し、さらに水で粘度 調製してウォッシュコート用スラリーを得た。このスラ リーを1.25 mmピッチのコージュライト社製のハニ カムに塗布して触媒を担持させ、試作サンブル (A-1 5)を得た。このときのスラリーの塗布量は、ハニカム 1 c c 当たり0.135g であった。

実施例16

実施例14において、シュウ酸パナジル水溶液に代えて メタタングステン酸アンモニウム水溶液(♥○。として 50重量%の水溶液)23.58gを加えたこと以外は 実施例14と同様にしてペロブスカイト化合物(La o. a Sro. 2 Coo. a Mo. 2 Os)を得た。 C のペロブスカイト化合物の比表面積は、13.6m2/ gであった。このようにして得たペロブスカイト化合物 30gと、住友化学社製のγ-アルミナ (商品名「A-11」) 100gとの混合物に、水を100g加え、遊 星ミルにて30分乾粉砕混合し、さらに水で粘度調製し てウォッシュコート用スラリーを得た。このスラリーを 1. 25 mmピッチのコージュライト社製のハニカムに 塗布して触媒を担持させ、試作サンブル(A-16)を 121g/1のNaOH水溶液を滴下し、pHを10と 50 得た。このときのスラリーの塗布量は、ハニカム1cc

11

当たり0.135gであった。 実施例17

La $(NO_3)_3 \cdot 6H_2 O$, Sr $(NO_3)_2$, Co (NO₃)₂・6H₂O、H₂PtCl₃・6H₂Oを 各々173.20g、126.98g、276.48 g、25.90gを秤量し1000mlの水に溶かし た。この水液に、121g/1のNaOH水溶液攪伴し ながら滴下し、pHを10とした。加水分解反応終了 後、18時間攪伴を続けた。以下実施例1と同様にし て、ペロプスカイト化合物(Lac Src Co 10 o. as Pto. os Os)を得た。このペロブスカイ ト化合物の比表面積は、27.6 m²/gであった。と のようにして得たペロブスカイト化合物30gと、住友 化学社製のγ-アルミナ(商品名「A-11」) 100 gとの混合物に、水を100g加え、遊星ミルにて30 分間粉砕混合し、さらに水で粘度調製してウォッシュコ ート用スラリーを得た。このスラリーを1.25mmビ ッチのコージュライト社製のハニカムに塗布して触媒を 担持させ、試作サンプル(A-17)を得た。このとき 3gであった。

実施例18

実施例17において、H2PtCle・6H2Oに代え TRh (NO₃) ₃ · 2H₂ Oを16. 25gを用いた こと以外は実施例17と同様の方法にして、ペロブスカ イト化合物(Lao. 4 Sro. e Coo. es Rh o. os Os)を得た。このペロブスカイト化合物の比 表面積は、29.6m~/gであった。このようにして 得たペロブスカイト化合物30gと、住友化学社製の7 -アルミナ(商品名「A-11」) 100gとの混合物 30 に、水を100g加え、遊星ミルにて30分間粉砕混合 し、さらに水で粘度調製してウォッシュコート用スラリ ーを得た。このスラリーを1.25mmピッチのコージ ュライト社製のハニカムに塗布して触媒を担持させ、試 作サンプル(A-18)を得た。このときのスラリーの 塗布量は、ハニカム1cc当たり0.129gであっ た。

実施例19

実施例17において、H₂PtCl。・6H₂Oに代え てPdCl。を8.67gを用いたこと以外は実施例1 7と同様にして、ペロプスカイト化合物 (Lao. 。S ro. a Coo、a s Pdo. o s Os)を得た。この ペロプスカイト化合物の比表面積は、28.5m2/g であった。このようにして得たペロプスカイト化合物3 0gと、住友化学社製のγ-アルミナ (商品名「A-1 1」) 100gとの混合物に、水を100g加え、遊星 ミルにて30分間粉砕混合し、さらに水で粘度調製して ウォッシュコート用スラリーを得た。このスラリーを 1. 25 mmピッチのコージュライト社製のハニカムに 塗布して触媒を担持させ、試作サンブル(A-19)を 50 実施例25

得た。このときのスラリーの塗布量は、ハニカム1 c c 当たり0.122gであった。

実施例20

実施例17において、H2PtCls・6H2Oに代え てRuCl₄・5H₂Oを16.15gを用いたこと以 外は実施例17と同様にしてペロブスカイト化合物(L ao. 4 Sro. 8 Coo. 8 5 Ruo. 0 5 O5) & 得た。このペロブスカイト化合物の比表面積は、25. 3 m² /gであった。このようにして得たペロブスカイ ト化合物30gと、住友化学社製のγ-アルミナ(商品 名「A-11」)100gとの混合物に、水を100g 加え、遊星ミルにて30分間粉砕混合し、さらに水で粘 度調製してウォッシュコート用スラリーを得た。このス ラリーを1.25mmピッチのコージュライト社製のハ ニカムに塗布して触媒を担持させ、試作サンブル(A-20)を得た。このときのスラリーの塗布量は、ハニカ ム1ccあたり0.129gであった。

実施例21

実施例3において、ペロブスカイト化合物(LaCoO のスラリーの塗布量は、ハニカム1cc当たり0.12 20 g)と、日本化学社製のH型モルデナイト(商品名「H M-23」) とを、各々1g、100g使用したこ以外 は実施例3と同様にして、1.25mmピッチのハニカ ム形状の試作サンプル(A-21)を得た。このときの スラリーの塗布量は、ハニカム1 c c 当たり0.929 gであった。

実施例22

実施例3において、ペロブスカイト化合物(LaCoO 。)と、日本化学社製のH型モルデナイト(商品名「H M-23」) とを、各々10g、100g使用したこと 以外は実施例3と同様にして、1.25mmピッチのハ ニカム形状の試作サンブル(A-21)を得た。このと きのスラリーの塗布量は、ハニカム1cc当たり0.9 48gであった。

実施例23

実施例3において、ペロブスカイト化合物(LaCoO 。)と、日本化学社製のH型モルデナイト(商品名「H M-23」) とを、各々50g、100g使用したこと 以外は実施例3と同様にして、1.25mmピッチのハ ニカム形状の試作サンブル(A-23)を得た。このと きのスラリーの塗布量は、ハニカム1cc当たり0.1 09gであった。

実施例24

実施例3において、ペロブスカイト化合物(LaCoO 。)と、日本化学社製のH型モルデナイト(商品名「H M-23」)とを、各々50g、100g使用したこと 以外は実施例3と同様にして、1.25mmピッチのハ ニカム形状の試作サンブル (A-24) を得た。このと きのスラリーの塗布量は、ハニカム1cc当たり0.1 22gであった。

組成式: Nax [(AlO₂)_x · (SiO₂)_y]· ZH2Oで表されるナトリウム型モルデナイトの市販品 (日本モービル社製、商品名「ZSM-5」、Y/X-35)100gを0.025モル/1のTiOSO4水 溶液 1 リットル中に浸漬し、充分に攪伴した。これをオ ートクレーブ中にて攪伴しながら100℃/時の昇温速 度で昇温して125℃に1時間保持し、TiOSO。を 加水分解させて、NaをTiでイオン交換した後、ろ 別、水洗してゼオライトのケーキを得た。次いで、この ケーキを乾燥した後、650℃で4時間焼成してゼオラ 10 イトを得た。このゼオライト中のTiの含有量はTiO 2 として、2. 4重量%であった。このようにして得た Ti-ZSM-5をH型モルデナイトに代えて用いたと と以外は実施例3と同様にして、1.25mmピッチの ハニカム形状の試作サンブル (A-25) を得た。この ときのスラリーの塗布量は、ハニカム1cc当たり0. 110gであった。

比較例1

実施例1において、活性酸化チタンを使用せず、ペロブ スカイト化合物 (Lao. 4 Sro. c Coo. g Mn 20 結果を表1~表3に示す。

。 20。) だけを用いてウォッシュコート用スラリー を得、その他は実施例1と同様にして、1.25mmピ ッチのハニカム形状の比較サンブル(B-1)を得た。 このときのスラリーの塗布量は、ハニカム1 c c 当たり 0.132gであった。

(2)評価試験

上記サンブル (A-1)~(A-25)並びに比較サン ブル(B-1)について、下記の試験条件でA/Fを1 4. 0~40の範囲を変化させることにより窒素酸化物 COおよびC。H。含有ガスの浄化試験を行った。

(試験条件)

(1)ガス組成 1000ppm NΟ

CO 1 %

C₃ H_e 1000ppm

10% CO2 A/F $14 \sim 40$

(2)空間速度 10000 1/Hr

(3) 反応温度 300℃, 400℃, 500

℃、又は600℃

15

表 1

反応温度300℃

16

					A/1	3			
		14.6			20.0			40.0	
	ΝО	нс	co	NO	нс	со	NO	нс	co
A-1	72	99	93	89	99	99	77	99	99
A-2	65	98	90	84	99	99	73	99	99
A-3	59	9 5	88	8 3	99	9 5	76	99	97
A-4	74	99	93	92	99	99	81	99	99
A-5	69	97	87	84	98	99	73	99	99
A-6	6 5	98	89	85	99	99	75	99	99
A – 7	84	99	99	95	99	99	87	99	99
A-8	74	97	96	86	99	99	76	88	99
A-9	71	95	9 0	86	98	93	78	99	98
A-10	79	99	93	73	88	99	65	99	99
A-11	68	98	9 5	8 1	99	98	72	99	99
A-12	65	98	9 2	84	99	99	75	99	88
A-13	70	99	95	92	99	99	81	99	9.9

		14.6		į	20.0			40.0	
	ON	нС	00	NO	нс	၀၁	NO	нс	00
A-14	6.2	9.7	88	80	9.7	9.1	7.3	98	95
A-15	ភភ	93	8.1	7.9	9 5	9.0	7.3	16	95
A-16	5.8	95	84	6 2	2 6	93	7 5	9 8	95
A-17	7.8	66	66	8 5	66	66	4.9	66	66
A-18	7.4	66	66	8 1	6 6	88	0.2	66	66
A-19	7.5	66	66	8 2	66	9.9	73	66	66
A-20	7.9	66	66	8.7	66	9.9	8 1	6 6	66
A-21	4 9	83	១១	56	7.1	62	45	74	8 8
A-22	7.5	9 8	9.2	7.0	66	98	61	66	66
A-23	7.9	66	93	7.3	66	9.9	65	6 6	6 6
A-24	8 2	66	66	65	66	99	58	6 6	66
A-25	6 1	9.7	06	85	66	98	7.8	6 6	6 6
B-1	7.0	66	9.2	4	66	66	0	66	6 6

(表1つらを)

19

A/F=20における温度特性

C)

••	3002			400°C			500°			200.9	
ON	нС	00	NO	HC	00	ON	HC	00	ON	нс	ပ
9.2	66	66	0.6	66	66	7.5	66	66	48	6 6	86

*表1~表2より、本発明に係る触媒〔試作サンブル(A-1)~(A-25)〕は、いずれも窒素酸化物、CO及びC。H。の浄化率が高いのに対して、比較触媒〔比較サンブル(B-1)および(B-2)〕は、総じてその浄化率が低い事がわかる。

20

【発明の効果】以上詳細に説明したように、本発明に係る三元触媒は、広いA/Fの範囲において排ガス中の窒素酸化物CO、および炭化水素を効率良く浄化することができるなど、本発明は優れた特有の効果を奏する。

10

20

30

* 40

フロントページの続き

	識別記号	庁内整理番号	FΙ	技術表示箇所
23/80	Α	8017-4G		
23/85	Α	8017-4G		·
23/89	Α	8017-4G		
29/04	Α	6750-4G		
29/18	Α	6750-4G		
	23/89 29/04	23/80 A 23/85 A 23/89 A 29/04 A	23/80 A 8017-4G 23/85 A 8017-4G 23/89 A 8017-4G 29/04 A 6750-4G	23/80 A 8017-4G 23/85 A 8017-4G 23/89 A 8017-4G 29/04 A 6750-4G

(12)

特開平5-76762

29/28

A 6750-4G



(11) Publication number:

05-076762

(43) Date of publication of application: 30.03.1993

(51)Int.CI.

B01J 23/84 B01D 53/36 B01J 23/76 B01J 23/78 B01J 23/80 B01J 23/85 B01J 23/89 B01J 29/18 B01J 29/18 B01J 29/18

(21)Application number: 03-315589

(22)Date of filing:

03-315589 21.09.1991 (71)Applicant:

SAKAI CHEM IND CO LTD

(72)Inventor:

NAKATSUJI TADAO OKUNO MASAO

YOSHIMOTO MASAFUMI

(54) CATALYST FOR CATALYTIC REDUCTION OF NITROGEN OXIDE

(57) Abstract:

PURPOSE: To increase the purification efficiency of a ternary catalyst for simultaneous removal of NOx, hydrocarbon compds. and CO in exhaust gas within a wide range of fuel-air ratio by supporting a perovskite type multiple oxide represented by a specified formula on a solid acid carrier.

CONSTITUTION: A perovskite type multiple oxide represented by a formula AxB1-xCyC1-yO3 is supported on zeolite or other solid acid carrier to obtain a catalyst for simultaneous removal of NOx, hydrocarbon compds. and CO in exhaust gas. In the formula, A is La or Ce, B is Ba, Sr, Ca, Mg, Pb, Zn or Co, C is Fe, Ni, Cr, Cu, V, Mo, W, Ti, Zr, Nb, Pd, Rh, Ru or Pt, $0 \le x \le 1$ and $0 \le y \le 1$. When the catalyst is used, NOx and hydrocarbon in exhaust gas are efficiently removed within a wide range of fuel-air ratio.

LEGAL STATUS

[Date of request for examination]

[Date of sending the examiner's decision of rejection]

[Kind of final disposal of application other than the examiner's decision of rejection or application converted registration]

[Date of final disposal for application]

[Patent number]

[Date of registration]

[Number of appeal against examiner's decision of rejection]

[Date of requesting appeal against examiner's decision of rejection]

[Date of extinction of right]

Copyright (C); 1998,2003 Japan Patent Office



CLAIMS DETAILED DESCRIPTION TECHNICAL FIELD EFFECT OF THE INVENTION TECHNICAL PROBLEM MEANS EXAMPLE

[Translation done.]

* NOTICES *

JPO and NCIPI are not responsible for any damages caused by the use of this translation.

- 1. This document has been translated by computer. So the translation may not reflect the original precisely.
- 2.**** shows the word which can not be translated.
- 3.In the drawings, any words are not translated.

CLAIMS

[Claim(s)]

[Claim 1] The catalyst for removing to coincidence the nitrogen oxides, hydrocarbon compound, and carbon monoxide in the exhaust gas characterized by solid acid support coming to support the perovskite mold multiple oxide expressed with following the (1) type. AXB1-XCYC1-YO3 (1)

(Among a formula, La, or Ce and B are [Mn, or Co and C of Ba, Sr, calcium, Mg Pb, Zn, or Ag and C] Fe, nickel, Cr, Cu, V, Mo, W, Ti, Zr, Nb, Pd, Rh, Ru, or Pt, and A is 0<=X<=1 and 0<=Y<=1.)

[Translation done.]

* NOTICES *

JPO and NCIPI are not responsible for any damages caused by the use of this translation.

1. This document has been translated by computer. So the translation may not reflect the original precisely.

2.**** shows the word which can not be translated.

3.In the drawings, any words are not translated.

DETAILED DESCRIPTION

[Detailed Description of the Invention]

[Industrial Application] This invention relates to the three way component catalyst used in case the harmful nitrogen oxides contained in the exhaust gas discharged from works, an automobile, etc., a hydrocarbon, and a carbon monoxide are removed to coincidence in detail with respect to the catalyst for nitrogen-oxides catalytic reduction.

[Description of the Prior Art] Since the reactions which remove injurious ingredients in exhaust gas, such as NOX, HC, and CO, to coincidence are the reaction which returns NOX to N2, and a reaction for which the reaction which oxidizes HC and CO in a carbon dioxide and water advances to coincidence and they become very important [the quantitative ratio of oxidizing quality components contained in exhaust gas, such as oxygen and NOX, and reducibility components such as HC and CO,], offgas treatment is performed on the conditions near theoretical air fuel ratio. However, operation by theoretical air fuel ratio became the cause of aggravation of fuel consumption, and development of a catalyst with a large window has been desired. Conventionally, as a three way component catalyst component, a platinum-rhodium-alloy, palladium-rhodium, and platinum-palladium-rhodium catalyst shows high activity, and the catalyst which supported the platinum rhodium alloy to the alumina is put in practical use. Moreover, since these precious metal catalysts are expensive, the catalyst which makes a perovskite mold multiple oxide or these contain palladium as a catalyst replaced with this has been proposed. however, any of these catalysts — although — only theoretical air fuel ratio or its neighborhood ratio showed high activity. (The width of face of a window is narrow) Therefore, development of the catalyst which has the window where width of face is wide has been desired, without sacrificing fuel consumption. This invention is made in view of the above situation, and the place made into the purpose is to offer the three way component catalyst which has the window where width of face is wide, and the three way component catalyst with which in other words the gap from theoretical air fuel ratio functions effectively on the bottom of a large exhaust gas condition.

[Means for Solving the Problem] Solid acid support, such as a zeolite, crystalline silicic acid aluminum phosphate (SAPO), crystalline aluminum phosphate (ALPO), crystalline phosphoric acid metal aluminum (MAPO), an alumina, a titania, a zirconia, and a silica alumina, comes to support the perovskite mold multiple oxide by which the catalyst for nitrogen-oxides catalytic reduction which uses the hydrocarbon and/or oxygenated compound concerning this invention as a reducing agent is expressed with following the (2) type. AXB1-XCYC1-YO3 (2)

the inside of a formula, and A — La, or Ce and B — Ba, Sr, calcium, Mg, Pb, Zn, or Ag — C is Co, and Mn or C is Fe, nickel, Cr, Cu, V, Mo, W, Ti, Zr, Nb, Pd, Rh, Ru, or Pt, and it is 0<=X<=1 and 0<=Y<=1. The solid acid support in this invention means the support which shows solid acid nature in the temperature field used. The check of solid acid nature is in using the temperature programmed desorption which used ammonia, ammonia, or a pyridine. situ It is made by the FTIR method. As solid acid support, there are zeolite system solid acid support, oxide system solid acid support, etc. which are shown below.

(i) Zeolite system solid acid support processes the zeolite which is excellent in thermal resistance, such as Na-mordenite, Na-ZSM -5, and Na-USY (USY: ultra stay bull Y mold zeolite), from acids, such as a water solution of ammonium salt, such as an ammonium sulfate, or a sulfuric acid, and is obtained by carrying out the ion exchange of some or all of alkali metal in a zeolite to ammonium ion (NH4+) or a hydrogen ion (H+). When based on the approach of carrying out the ion exchange by NH4+, finally baking processing is needed. It is the acid type mordenite obtained by carrying out acid treatment of the mordenite mold zeolite expressed with following the (3) type, for example as zeolite system solid acid support. The acid type mordenite whose mole ratios of SiO2/aluminum 2O3 are 13-20, and are 25-200, The zeolite obtained by carrying out the ion exchange of some or all of Ion M in the zeolite expressed with following the (4) type by Ti4+, Zr4+, or Sn4+ is mentioned.

M2 [(AIO2) two r(SiO2) 10] -ZH2O (3)

(However, it is the value to which M is changed with alkali-metal ion among a formula, and r is changed according to the synthetic conditions of a zeolite.)

M'A [(AlO2) p(SiO2) a] -Z'H2O (4)

(However, ion M' is alkali-metal ion, alkaline-earth-metal ion or a hydrogen ion, nA=p (n is the valence of Ion M), and q/p>=5 among a formula.)

(ii) — as oxide system solid acid support — aluminum2 — 03, TiO2, TiO2/SO4, —, ZrO2, and ZrO2/SO4 — single metallic oxides, such as —, the multiple oxide of SiO2/aluminum 2O3, TiO2/aluminum 2O3, and TiO2/ZrO2 grade, etc. are mentioned. In these, aluminum 2O3 and SiO2 [ZrO2 and]/aluminum 2O3 are desirable in respect of thermal resistance.

(iii) The crystalline aluminum phosphate (ALPO) of a kind which has the porous structure or the layer structure of zeolite resemblance as other examples of solid acid support, the crystalline phosphoric acid metal aluminum (MAPO) which permuted a part of crystalline silicic acid aluminum phosphate (SAPO) which is the close relationship matter, phosphorus of ALPO, or phosphorus—aluminum with metals, such as titanium, iron, magnesium, zinc, manganese, and cobalt, are mentioned. ALPO type phosphate can be adjusted to the combination of the request chosen from the above—mentioned phosphagen and the source of a metal, a silica and a silica sol, silicate of soda, etc. with a hydrothermal crystallization method under the case where a zeolite is compounded, and similar conditions, from the

raw material which mixed the so-called templates, such as an amine and quaternary ammonium. The main differences with the case where a zeolite is compounded are generally compounded more in pH acidity field by the elevated temperature (in general 150 degrees C or more). Generally the presentation of ALPO type phosphate is aluminum 203. -(0.8-1.2)— It is expressed with P2O5 and nH2O. Moreover, although the peaks of the silica which is permuted in SAPO or MAPO, and a metal are about about 1 of the total amount of aluminum and phosphorus / 10, they may use in this invention what is not necessarily contained in this presentation range, i.e., the thing containing an amorphous substance. When using as support the phosphate of the ALPO type obtained by the hydrothermal crystallization method, after rinsing and drying, generally what carried out incineration removal of the template which calcinates and remains in air is used. The catalyst concerning this invention can be adjusted by the approach of of (1), (2), or (3) shown below. In the slurry which distributed solid acid support, (1) La, Ce, Ba, Sr, calcium, Mg, Pb, Zn, Ag, Mn, Co, Fe, nickel, Cr, Cu, By water—soluble salts, such as a nitrate of V, Mo, W, Ti, Zr, Nb, Pd, Rh, Ru, or Pt, the method of throwing an alcoholic solution into these alkoxides and making these neutralize or hydrolyze, etc. Solid acid support is made to support perovskite compound precursors, such as these compound metal kinds of hydroxide. Subsequently, it dries and calcinates, after repeating filtration, rinsing, and repulping and performing them.

(2) Fully carry out wet-grinding mixing of solid acid support and the perovskite compound adjusted separately by a planetary mill etc. (3) Precursors, such as a water-soluble salt of solid acid support, or a hydroxide, and La, Ce, Ba, Sr, calcium, Mg, Pb, Zn, Ag, Mn, Co, Fe, nickel, Settlings are made to generate by water-soluble salts, such as a nitrate of Cr, Cu, V, Mo, W, Ti, Zr, Nb, Pd, Rh, Ru, or Pt, the method of making the solution which mixed the alcoholic solution of these alkoxides to homogeneity neutralize or hydrolyze, etc. Subsequently, it dries and calcinates, after repeating filtration, rinsing, and repulping and performing these settlings. In the above approach, the lower one of the generation temperature of a perovskite compound is desirable. The reason is that it is avoidable that the perovskite compound which has a big specific surface area is obtained, and the solid acid nature of solid acid support deteriorates by the reaction of solid acid support and the element which constitutes a perovskite compound, or the activity of a catalyst falls by the fall of the amount of generation of a perovskite compound, so that the generation temperature is low. therefore, aluminum2 — if it is when using solid acid support with high reactivity with the element which constitutes a perovskite compound like O3 and TiO2, the approach of of (3) which raises the homogeneity of the element which constitutes solid acid support, and the element which constitutes a perovskite compound is not desirable. Generally, although the approach of (1) is desirable, the catalyst which shows quite high activity also by the approach of (2) can be acquired. The suitable amount of support of a perovskite compound is 1.0 - 50 % of the weight to the AUW of this perovskite compound and solid acid support. If 50 % of the weight is exceeded, although the rate of purification of a hydrocarbon or a carbon monoxide by oxygen becomes high in the system of reaction with which the addition effectiveness according to increase in quantity is not not only acquired, but oxygen coexists, the rate of purification of NOX will fall greatly. On the other hand, at less than 1.0 % of the weight, NO, a hydrocarbon, and any rate of purification of CO cannot be raised enough. The catalyst concerning this invention can be conventionally fabricated in the shape of a honeycomb, and the various configurations of spherical ** by the well-known shaping approach. In the case of this shaping, a shaping assistant, a Plastic solid reinforcement object, an inorganic fiber, an organic binder, etc. may be blended suitably. Moreover, covering support can be carried out by the wash coat method etc. on the base material fabricated beforehand. In addition, it can also be conventionally based on the wellknown catalyst preparation. Although the optimal temperature the catalyst concerning this invention indicates purification activity to be to nitrogen oxides, a hydrocarbon, and a carbon monoxide changes with catalyst kinds, it is usually 100-800 degrees C, and it is desirable in this temperature field to carry out conduction of the exhaust gas about (space-velocity SV) 500-100,000. In addition, a more suitable service temperature field is 200-800 degrees C. moreover, the air-fuel ratio (A/F) as which the catalyst concerning this invention functions effectively — 14.0-60 — it is — desirable — 14.0-40 — it is 14.0-30 more preferably.

[Example] It is possible to change this invention suitably in the range which is not limited to the following example at all and does not change the summary, and to carry out hereafter, although this invention is further explained to a detail based on an example. (1) Respectively, 101.05g, 135.83g weighing capacity of adjustment example 1La(NO3) 2.6H2O of a catalyst, Mn(Ac)2.4H2O (the same is said of Ac:CH3COOH and the following), Sr (NO3)2, and the Co(NO3)2.6H2O was carried out, and they were melted in 500ml water 74.08g 28.60g. The NaOH water solution of 121 g/l was added to this solution, fully stirring, and pH was set to 10. It riped by continuing stirring for 18 hours after neutralization termination. then, filtration, rinsing, and repulping — the conductivity of filtered water repulping — it repeated until it became almost the same as service water. The obtained filter cake was dried at 120 degrees C for 18 hours, and, subsequently it calcinated at 700 degrees C for 3 hours. As a result of calculating XRD of the obtained baking object, it turned out that the perovskite crystal phase is generating. Moreover, the specific surface area (the following specific surface area is also depended on the law) by the BET adsorption method of this baking object was 23.7m2/g (La0.4Sr0.6Co0.8Mn 0.2O3). Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of activity titanium oxide which calcinated the metatitanic acid (TiO2andH2O) obtained according to the sulfuric-acid method titanium oxide process for 3 hours, and obtained it at 600 degrees C] (specific surface area: 104.2m2/g) mixture, in the planetary mill, grinding mixing was carried out for 30 minutes, viscosity control was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-1) was obtained. The coverage of the slurry at this time was 0.116g per honeycomb 1cc.

50.66g weighing capacity of example 2La(NO3)2.6H2O and the 89.5g Mn(Ac)2.4H2O was carried out respectively, and, subsequently the perovskite compound (LaMnO3) was obtained by the same approach as Example I. The specific surface area of this perovskite compound was 29.1m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-2) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.094g per honeycomb 1cc.

50.66g weighing capacity of example 3La(NO3)2.6H2O, Pb (NO3)2, and the 71.60g Mn(Ac)2.4H2O [13.69g] was carried out respectively, it mixed, and, subsequently the perovskite compound (La0.8Pb0.2MnO3) was obtained by the same approach as Example I except having calcinated at 800 degrees C for 3 hours. The specific surface area of this perovskite compound was 23.7m2/g. Thus, in

addition to [100g of water] 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-3) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.117g per honeycomb 1cc.

59.19g weighing capacity of example 4La(NO3) 2.6H2O and the 88.07g Co(NO3)2.6H2O was carried out respectively, it mixed, and, subsequently the perovskite compound (LaCoO3) was obtained by the same approach as Example I except having calcinated at 800 degrees C at 3:00. The specific surface area of this perovskite compound was 17.47m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and H form mordenite (HM-23) 100g [by the Japanization study company] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-4) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.113g per honeycomb 1cc.

59.19g weighing capacity of example 5Ce(NO3)2.6H2O, Ba (NO3)2, and the 70.65g Co(NO3)2.6H2O [10.63g] was carried out respectively, and, subsequently the perovskite compound (Co0.8Ba0.2CoO3) was obtained by the same approach as an example 1. The specific surface area of this perovskite compound was 23.0m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and H form mordenite (trade name "HM-23") 100g [by the Japanization study company] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-5) of 1.25mm pitch honeycomb configuration was obtained. The coverage of the slurry at this time was 0.130g per honeycomb 1cc. 339.0ml weighing capacity of example 6La(NO3) 2.6H2O, Mn(Ac)2.4H2O, and the 90.84g (water solution of 14.82g/100ml concentration as Ti) of the 25.71g of the TiCl4 water solutions was carried out respectively, and, subsequently the perovskite compound (LaMn0.5Ti 0.5O3) was obtained by the same approach as Example I. The specific surface area of this perovskite compound was 25.3m2/g. Thus, in addition to [100g of water] 10g of obtained perovskite compounds, and silica-alumina (trade name "COK-84") 100g [made from Japanese AROJIRU] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-6) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.098g per honeycomb 1cc.

The ethanol solution of La ethoxide by example 7 Freparation of perovskite compound
Hakusui Chemical Industries (solution of 73 g/l concentration as La 2O3), 9.22ml weighing capacity is carried out and it mixes, and stirring enough, 1% of the weight of the aqueous ammonia solution was dropped gradually, and the ethanol solution (solution of 79 g/l concentration as BaO) of company Ba ethoxide, the ethanol solution (solution of 67 g/l concentration as NiO) of company nickel ethoxide, and 49.95ml (solution of 91g [/l.] concentration as CoO) of 21.74ml of 100.0ml of ethanol solutions of Co ethoxide were made to hydrolyze respectively. Subsequently, evaporation to dryness was carried out, carrying out homogeneity mixing of this, it calcinated at 600 degrees C for 3 hours, and the perovskite compound (La0.8Ba0.2Coo.8nickel 0.2O3) was obtained. The specific surface area of this perovskite compound was 36.9m2/g.

Aluminum isopropoxide 90.7g broken finely was added to 129.6g of of SAPO-34> water small quantity every, stirring, and stirring mixing was carried out until it became homogeneity. 51.3g of phosphoric acid water solutions was dropped at this mixed liquor 85%, and after carrying out stirring mixing until it became homogeneity, stirring mixing was fully carried out [silica sol 16.0g] 50 more%. Subsequently, hydroxylation tetraethylammonium 81.6g was added and stirring mixing was fully carried out. After having carried out filtration separation of the product after it taught this mixture to the autoclave and it carried out the stirring reaction at 200 degrees C for 24 hours, and rinsing and drying further, it calcinated in 3-hour air at 500 degrees C, and SAPO-34 were obtained. These SAPO-34 were the thing of 9.5 and the presentation contained 18.0 or 19.0% of the weight about Si, aluminum, and P, respectively. Thus, 100g of water was added to the mixture of perovskite compounds [which were obtained / 25g and 100g] SAPO-34, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A–7) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.138g per honeycomb 1cc. Weighing capacity of 101.05g, 74.08g, 135.83g, and the 33.60g was carried out respectively, example 8La(NO3)3.6H2O, Sr (NO3)2, Co (NO3) 2.6H2O, and Fe(NO3)2.6H2O were mixed, and the perovskite compound (La0.4Sr0.6Co0.8Fe 0.2O3) was obtained by the same approach as an example 1. The specific surface area of this perovskite compound was 21.6m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and 100g [of gamma–alumina by Sumitomo Chemical Co., Ltd.] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A−8) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.098g per honeycomb

Weighing capacity of 202.10g, 28.41g, 135.83g, and the 28.19g was carried out respectively, example 9La(NO3) 3.6H2O, Zn(NO3) 2.6H2O, Co(NO3) 2.6H2O, and Cu(NO3)2 and 3H2O were mixed, and the perovskite compound (La0.8Zn0.2Co0.8Cu 0.2O3) was obtained by the same approach as an example 1. The specific surface area of this perovskite compound was 17.3m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-9) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.083g per honeycomb 1cc.

Weighing capacity of 202.10g, 19.82g, 135.83g, and the 69.87g was carried out respectively, example 10La(NO3)3.6H2O, AgNO3, Co (NO3) 2.6H2O, and Zr(NO3) 4 and 5H2O were mixed, and the perovskite compound (La0.8Ag0.2Co0.8Zr 0.2O3) was obtained by the same approach as an example 1. The specific surface area of this perovskite compound was 17.3m2/g. Thus, in addition to [100g of water] 30g of obtained perovskite compounds, and 100g [of gamma—alumina by Sumitomo Chemical Co., Ltd.] mixture, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-10) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.097g per honeycomb 1cc.

example 11La(NO3)3.6H2 — O, Sr (NO3)2, and Co(NO3)2.6H2 — weighing capacity of each 101.05g, 74.08g, 135.83g, and 27.79g was

carried out, O and Cr (NO3)3 were mixed, and the perovskite compound (La0.4Sr0.6Co0.8Cr o.2O3) was obtained by the same approach as an example 1. The specific surface area of this perovskite compound was 20.3m2/g.

45.8g (what contains 9.5% of acetic acids alumina 67%) of pseudo-boehmite powder was added to 69.2g of phosphoric acid, and 178g of water] mixture small quantity every preparation of ALPO-5> 85%, and stirring mixing was carried out until it became homogeneity. Tripropylamine 43.8g was added to this liquid, and stirring mixing was carried out until it became homogeneity. After teaching this mixture to the autoclave and carrying out a stirring reaction at 150 degrees C for 70 hours, filtration separation of the product was carried out, after rinsing and drying, it calcinated in 3-hour air at 500 degrees C, and ALPO-5 were obtained. These ALPO-5 were the thing of the presentation which contains aluminum and P 18.0 or 22.0% of the weight, respectively. Thus, in addition to [100g of water] the mixture of perovskite compounds [which were obtained / 30g and 100g] ALPO-5, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-11) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.104g per honeycomb 1cc. 216.81ml weighing capacity of the 74.08g 101.05g [135.83g] was carried out respectively, it mixed, example 12 ⟨preparation of perovskite compound> La(NO3)2.6H2O, Sr (NO3)2, Co(NO3)2.6H2O, and NbCl5 (a hydrochloric-acid water solution, solution of 50 g/l concentration as Nb) were made into the example 1 below, and the perovskite compound (La0.4Sr0.6Co0.8Nb 0.2O3) was obtained. The specific surface area of this perovskite compound was 18.9m2/g.

Stirring mixing was carried out until it added aluminum iso prop KISHIDO 56.3g broken finely small quantity every and became homogeneity, stirring first manganese of preparation of MAPO-5> acetic acid 4.9g, and 4.1g of cupric acetate in the liquid which dissolved in 129g of water. Stirring mixing was carried out until it became homogeneity in this liquid [small quantity every], stirring 55.4g [of phosphoric acid], and diethyl ethanolamine 56.3g, and the mixed liquor of 55.5g of water 85%. After teaching this liquid to the autoclave and making it react at 200 degrees C for 25 hours, filtration separation of the product was carried out, after rinsing and drying, it calcinated in 3-hour air at 500 degrees C, and MAPO-5 were obtained. These MAPO-5 were the thing of 19.0, 19.0, and the presentation contained 2.8 or 4.4% of the weight about aluminum, P. Mn, and Cu, respectively. Thus, in addition to [100g of water] the mixture of perovskite compounds [which were obtained / 25g and 100g] MAPO-5, the slurry was obtained, subsequently applied to the same honeycomb as an example 1, the catalyst was made to support, and the prototype sample (A-12) of the honeycomb configuration of 1.25mm pitch was obtained. The coverage of the slurry at this time was 0.116g per honeycomb 1cc. In example 13 example 1, it replaced with activity titanium oxide and the prototype sample (A-13) was obtained like the example 1 except having used ZrO2 (specific surface area: 148.3m2/g) which calcinated zirconium hydroxide for 3 hours and obtained it at 600

degrees C. The coverage of the slurry at this time was 0.139g per honeycomb 1cc.

Respectively, 88.07g, 50.66g weighing capacity of example 14 \(\sqrt{preparation}\) of perovskite compound
\(\sum{La(NO3)2.6H2O}\). Sr (NO3)2, and the Co(Ac)2.4H2O was carried out, and they were melted in 500ml water 10.76g. It was dropped fully stirring the NaOH water solution of 121 g/l, and pH of liquid was set to 10. It riped by continuing stirring for 18 hours after neutralization termination, then, filtration, rinsing, and RIPARUBU — the conductivity of filtered water — RIPARUBU — after repeating until it became almost the same as it of service water, the filter cake was dried at 120 degrees C for 18 hours. this dry matter was ground, 25.90ml (water solution of 100 g/l concentration as V) of oxalic acid vanadyl solutions was added to this grinding object, and it kneaded enough — evaporation to dryness was carried out, and it dried at 120 degrees C for 18 hours, subsequently it calcinated at 850 degrees C for 3 hours, and the perovskite compound (La0.8Sr0.2Co0.8V0.2O3) was obtained. The specific surface area of this perovskite compound was 12.8m2/g. It mixed enough, having carried out 97.20g weighing capacity of the 100.0g of the silica sol O molds (thing of 20-% of the weight concentration as SiO2) and zirconium chlorides (ZrCl4) by the preparation > Nissan chemistry company of a <silica-zirconia, and stirring them respectively, and the total amount was set to 500ml with water. The NaOH water solution of 121 g/l was dropped at this liquid, and pH was set to 10. Stirring was continued after neutralization termination for 18 hours, filtration, rinsing, and RIPARUBU were repeated after that, the filter cake was obtained, at 120 degrees C, it dried for 18 hours and this filter cake was calcinated for 3 hours. The specific surface area of the obtained baking object was 297m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of the above-mentioned baking objects] mixture, ***** mixing during 30 minutes was carried out in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-14) was obtained. The coverage of the slurry at this time was 0.127g per honeycomb 1cc.

In the example 15 Freparation of perovskite compound example 14, the perovskite compound (La0.8Sr0.2Co0.8Mo 0.2O3) was obtained like the example 14 except having replaced with the oxalic acid vanadyl water solution, and having added 260.22ml (water solution of 25 g/l concentration as MoO2) of ammonia nature water solutions of an ammonium molybdate.

It held at 70 degrees C for 1 hour, having been immersed in the zirconium nitrate water solution (referred to as ZrO2 thing of 100 g/I concentration), and stirring Na mordenite (NM-100P) 100g by the ⟨preparation of Zr-mordenite⟩ Japanization study company, and the ion exchange of Na was carried out to Zr. It calcinated at 650 degrees C after drying the zeolite cake rinsed [was filtered and] and obtained for 4 hours. The content of Zr of this zeolite (Zr-mordenite) was 3.3 % of the weight, and specific surface area was 391m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and Zr-mordenite 100g mixture, grinding mixing was carried out for 30 minutes in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-15) was obtained. The coverage of the slurry at this time was 0.135g per honeycomb 1cc.

In example 16 example 14, the perovskite compound (La0.8Sr0.2Co0.8M0.2O3) was obtained like the example 14 except having replaced with the oxalic acid vanadyl water solution, and having added 23.58g (50% of the weight of water solution as WO3) of ammonium metatungstate water solutions. The specific surface area of this perovskite compound was 13.6m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] (trade name "A-11") mixture, ***** mixing was carried out in the planetary mill for 30 minutes, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-16) was obtained. The coverage of the slurry at this time was 0.135g per honeycomb



Respectively, weighing capacity of 173.20g, 126.98g, 276.48g, and the 25.90g was carried out, and example 17La(NO3)3.6H2O, Sr (NO3) 2, Co(NO3)2.6H2O, and H2PtCl6.6H2O were melted in 1000ml water. While 121 g/l carried out NaOH water—solution stirring, it was dropped at this water, and pH was set to 10. Stirring was continued after hydrolysis reaction termination for 18 hours. The perovskite compound (La0.4Sr0.6Co0.95Pt 0.05O3) was obtained like the example 1 below. The specific surface area of this perovskite compound was 27.6m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] (trade name "A-11") mixture, grinding mixing was carried out for 30 minutes in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-17) was obtained. The coverage of the slurry at this time was 0.123g per honeycomb 1cc.

In example 18 example 17, it replaced with H2PtCl6.6H2O, and Rh(NO3)3.2H2O was made into the same approach as an example 17 except having used 16.25g, and the perovskite compound (La0.4Sr0.6Co0.95Rh 0.05O3) was obtained. The specific surface area of this perovskite compound was 29.6m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of gamma—alumina by Sumitomo Chemical Co., Ltd.] (trade name "A-11") mixture, grinding mixing was carried out for 30 minutes in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-18) was obtained. The coverage of the slurry at this time was 0.129g per honeycomb 1 cc.

In example 19 example 17, the perovskite compound (La0.4Sr0.6Co0.95Pd 0.05O3) was obtained like the example 17 except having replaced with H2PtCl6.6H2O, and having used 8.67g for PdCl2. The specific surface area of this perovskite compound was 28.5m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] (trade name "A-11") mixture, grinding mixing was carried out for 30 minutes in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-19) was obtained. The coverage of the slurry at this time was 0.122g per honeycomb 1cc.

In example 20 example 17, the perovskite compound (La0.4Sr0.6Co0.95Ru 0.05O3) was obtained like the example 17 except having replaced with H2PtCl6.6H2O, and having used 16.15g for RuCl4 and 5H2O. The specific surface area of this perovskite compound was 25.3m2/g. Thus, 100g of water was added to 30g of obtained perovskite compounds, and 100g [of gamma-alumina by Sumitomo Chemical Co., Ltd.] (trade name "A-11") mixture, grinding mixing was carried out for 30 minutes in the planetary mill, viscosity preparation was carried out further with water, and the slurry for wash coats was obtained. Applied this slurry to the honeycomb by the KOJU light company of 1.25mm pitch, the catalyst was made to support, and the prototype sample (A-20) was obtained. The coverage of the slurry at this time was 0.129g per honeycomb 1cc.

In example 21 example 3, the prototype sample (A-21) of the honeycomb configuration of 1.25mm pitch was obtained like the example 3 except ** which used a perovskite compound (LaCoO3) and 1g (trade name "HM-23") of 100g of H mold mordenites by the Japanization study company respectively. The coverage of the slurry at this time was 0.929g per honeycomb 1cc.

In example 22 example 3, the prototype sample (A-21) of the honeycomb configuration of 1.25mm pitch was obtained like the example 3 except having used a perovskite compound (LaCoO3) and 10g (trade name "HM-23") of 100g of H mold mordenites by the Japanization study company respectively. The coverage of the slurry at this time was 0.948g per honeycomb 1cc.

In example 23 example 3, the prototype sample (A-23) of the honeycomb configuration of 1.25mm pitch was obtained like the example 3 except having used a perovskite compound (LaCoO3) and 50g (trade name "HM-23") of 100g of H mold mordenites by the Japanization study company respectively. The coverage of the slurry at this time was 0.109g per honeycomb 1cc.

In example 24 example 3, the prototype sample (A-24) of the honeycomb configuration of 1.25mm pitch was obtained like the example 3 except having used a perovskite compound (LaCoO3) and 50g (trade name "HM-23") of 100g of H mold mordenites by the Japanization study company respectively. The coverage of the slurry at this time was 0.122g per honeycomb 1cc.

Example 25 empirical formula: It was immersed into 1I. of 0.025 mols [/l.] TiOSO4 water solutions, and 100g (made in Japanese Mobile, a trade name "ZSM-5", Y/X -35) of commercial items of the sodium mold mordenite expressed with Nax[(AlO2) Xand(SiO2) Y], and ZH2O was fully stirred. After having carried out the temperature up with the programming rate of 100 degrees C/o'clock, having held at 125 degrees C for 1 hour, making TiOSO4 hydrolyze and carrying out the ion exchange of Na by Ti, stirring this in an autoclave, the ** exception, it rinsed and the cake of a zeolite was obtained. Subsequently, after drying this cake, it calcinated at 650 degrees C for 4 hours, and the zeolite was obtained. The content of Ti in this zeolite was 2.4 % of the weight as TiO2. Thus, the prototype sample (A-25) of the honeycomb configuration of 1.25mm pitch was obtained like the example 3 except having replaced with and used obtained Ti-ZSM -5 for H mold mordenite. The coverage of the slurry at this time was 0.110g per honeycomb 1cc.

In example of comparison 1 example 1, activity titanium oxide was not used, but the slurry for wash coats was obtained only using the perovskite compound (La0.4Sr0.6Co0.8Mn 0.2O3), and others obtained the comparison sample (B-1) of the honeycomb configuration of 1.25mm pitch like the example 1. The coverage of the slurry at this time was 0.132g per honeycomb 1cc.

(2) The purification trial of nitrogen oxides CO and C3H6 content gas was performed by A/F changing the range of 14.0-40 to the evaluation test above-mentioned sample (A-1) - (A-25) a list by the following test condition about a comparison sample (B-1). (Test condition)

(1) Gas presentation NO 1000ppmCO 1%C3H6 1000ppmCO2 10%A/F 14-40 (2) space velocity 10000 I/Hr (3) reaction temperature 300 degrees C, 400 degrees C, 500 degrees C, or 600-degree-C result is shown in Table 1 - 3.

表]



反応温度300℃

					A / 1	?			
		14.6			20.0			40.0	
	NO	нс	co	NO	нс	co	NO	нс	co
A-1	72	99	93	89	99	99	77	99	99
A-2	65	98	9 0	84	99	99	73	9 9	99
A - 3	59	95	88	8 3	99	95	76	99	97
A - 4	74	88	93	92	99	99	81	99	99
A – 5	69	97	87	84	98	99	73	99	99
A-6	65	98	89	8 5	99	99	75	99	99
A-7	84	99	99	9 5	99	99	87	99	99
A-8	74	97	96	8 6	9 9	99	7 6	99	99
A-9	71	95	90	8 6	98	93	78	99	98
A-10	79	99	93	73	99	99	65	99	99
A-11	68	98	9 5	81	99	98	72	99	99
A-12	65	98	9 2	8 4	99	99	75	99	99
A-13	70	99	95	92	88	99	81	99	99

(表1つが巻)

											· -]			
	CO	9 5	95	95	66	66	66	6 6	68	66	66	66	66	66
40.0	нс	9 8	9.7	98	66	66	66	9 9	74	66	66	6 6	6 6	66
	ON	7.3	7.3	2 2	6 L	10	73	8 1	45	6.1	9	2.8	7.8	0
	00	9 1	06	93	66	9.8	66	6 6	6.2	98	66	66	8 6	66
20.0	нС	9.7	9 5	26	66	66	66	66	7.1	66	66	66	66	86
	ON	80	4.9	4 9	85	8 1	8 2	8 7	99	10	7.3	65	8 5	4
	.02	88	8 1	84	6 6	66	9.9	6 6	5 5	9.2	93	66	06	9.2
14.6	HC	9.7	8 8	95	66	66	66	66	63	98	66	66	9.7	66
	ON	62	ភភ	58	7.8	7.4	7.5	7.9	4 9	7.5	7.9	8 2	6.1	7.0
		A-14	A-15	A-16	A-17	A-18	A-19	A-20	A-21	A-22	A-23	A-24	A-25	B-1

LL	
型	
文	
関	
펦	
Ŕ	
*	
华	
77	
0	
ମ	
11	
14	
1	
⋖	

O

	00	6 6
600°C	нС	6 6
	NO	4 8
	00	66
500c	нс	66
	ON	75
	00	6 6
400°C	HC	G G
	ON	0 6
	00	9.6
3008	HC	66
	ON	9.2

From Table 1 – 2, the catalyst [prototype sample (A-1) concerning this invention – (A-25)] all understand that the rate of purification is low generally to a thing with high nitrogen oxides, CO, and rate of purification of C3H6 for a comparison catalyst [comparison sample (B-1) and (B-2)].

[Effect of the Invention] As explained to the detail above, this invention does the outstanding characteristic effectiveness so — the three way component catalyst concerning this invention can purify efficiently the nitrogen oxides CO and the hydrocarbon in exhaust gas in the range of large A/F.

[Translation done.]

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:

□ BLACK BORDERS
□ IMAGE CUT OFF AT TOP, BOTTOM OR SIDES
□ FADED TEXT OR DRAWING
□ BLURRED OR ILLEGIBLE TEXT OR DRAWING
□ SKEWED/SLANTED IMAGES
□ COLOR OR BLACK AND WHITE PHOTOGRAPHS
□ GRAY SCALE DOCUMENTS
□ LINES OR MARKS ON ORIGINAL DOCUMENT
□ REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY

IMAGES ARE BEST AVAILABLE COPY.

☐ OTHER:

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.